S/076/62/036/008/010/011 B101/B144

AUTHOR:

Zharkova, L. A.

TITLE:

Determination of thermodynamic functions for titanates and

silicates by comparative calculation

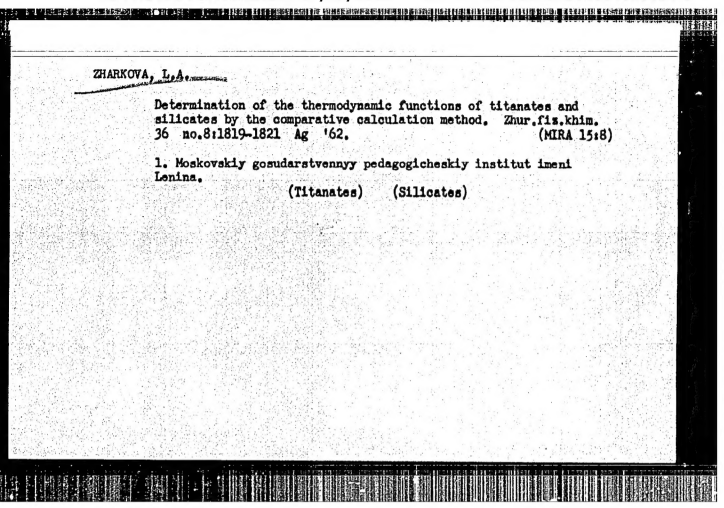
PERIODICAL: Zhurnal fizicheskoy khimii, v. 36, no. 8, 1962, 1819 - 1821

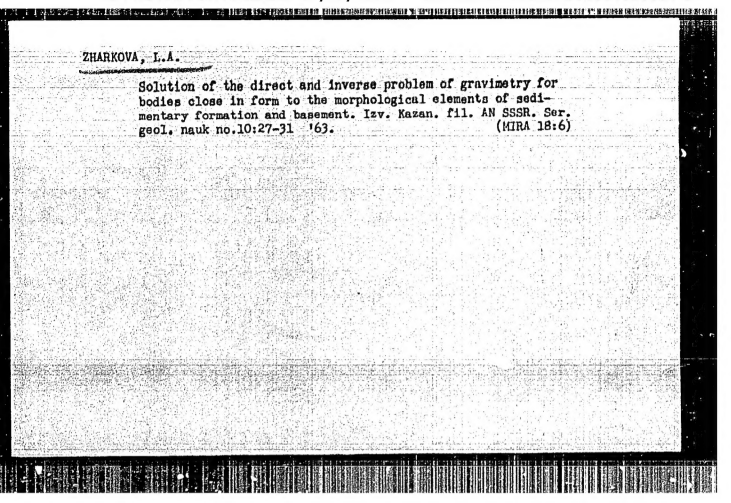
TEXT: Thermodynamic data for carbonates, meta- and orthosilicates, meta- and orthotitanates were calculated on the basis of equations suggested by M. Kh. Karapet'yants (Zh. fiz. khimii, 28, 353, 1954) establishing a relationship between the thermodynamic functions of these compounds, and were compared with published data. For carbonates, metasilicates, and metatitanates:  $\Delta Z_{298}^{0} = 0.993 \Delta H_{298}^{0} + 17.92$  with an accuracy of  $\pm 0.6$  kcal/mole. For orthotitanates:  $\Delta Z_{298}^{0} = 0.996 \Delta H_{298}^{0} + 25.3$ . The following relation exists between the thermodynamic functions of metatitanates and carbonates:  $\Delta Z_{298}^{0} = 1.002 \Delta Z_{298}^{0} = 1.002 \Delta Z_{298}^{0} = 1.003 \Delta Z_{298}^{0}$ 

Card 1/3

\$/076/62/036/008/010/011 Determination of thermodynamic ...  $\Delta z_{298}^{\circ} (\Delta H_{298}^{\circ}) = 1.001 \Delta z_{298}^{\circ} (\Delta H_{298}^{\circ}) - 90.8; \Delta z_{298}^{\circ} (\Delta H_{298}^{\circ}) = 1.001 \Delta z_{298}^{\circ}$ MeSiO3 - 16.8. For calculating the functions of orthotitanates and orthosilicates the following is suggested:  $\Delta Z_{298}^{\text{C}/\text{H}_{298}} = 1.01 \Delta Z_{298}^{\text{C}/\text{H}_{298}} - 24.2. \Delta Z_{298}^{\text{C}/\text{H}_{298}}$ Me Tio and 2 H<sub>298</sub> were calculated for 23 titanates and silicates not yet studied by experiment (Table 4). There are 3 figures and 4 tables. ASSOCIATION: Moskovskiy gosudarstvennyy pedagogicheskiy institut im. V. I. Lenina (Moscow State Pedagogical Institute imeni V. I. Lenin) January 6, 1962 SUBMIT PED: Table 4. Thermodynamic data of some meta- and orthotitanates, meta- and orthosilicates not yet studied by experiment. Legend: (1) compound; (2) kcal/mole. Card 2/3

<u> </u>	ΔH 238. ΚΚΩΑ/ΜΟΛΟ D	∆2°208. жкал/молцД	Соединение	ΔH <sub>208</sub> , жкал/молыΩ	AZ <sub>208</sub>	
Liatio, Naitio, Naitio, Katio, Csitio, Zntio, Cutio, Nitio, Cotio, Mntio, Phtio, Pbio,	376,0 3	-377, 2 -356, 0 -359, 6 -353, 1 -281, 2 -230, 2 -253, 2 -259, 0 -301, 8 -256, 1 -347, 4 -312, 7	Lisios Kasios Casios Srsios Basios Cusios Fesios Cosios Nisios Srasios	-381,5 -365,0 -355,5 -380,8 -381,4 -233,1 -260,6 -267,2 -516,3 -508,7	-360,8 -342,5 -336,0 -362,3 -363,3 -214,7 -252,1 -243,6 -257,3 Tab -488,8 -482,0	ıle 4





出租 表现 无法确定条款的可能是型。因的使用压制,我还通过医验性剂清洁是非自己的结婚,此时的生物的企用。由 66183 807/20-128-5-37/67 5.4700 Zharkova, L. A., Gerasimov, Ya. I., Corresponding Member, AS USSR, AUTHORS: Rezukhina, T. N., Simanov, Yu. P. The Equilibrium Between Zinc Tungstate and Hydrogen and the TITLE: Thermodynamic Characteristics of ZnWOA Doklady Akademii nauk SSSR, 1959, Vol 128, Nr 5, pp 992-994 PERIODICAL: (USSR) The thermodynamic characteristics of tungstates and molybdates of bivalent metals have been investigated at the laboratory of the ABSTRACT: authors since 1944. The method of circulation applied so far was not applicable here as zinc evaporates and is carried away from the reaction zone. The method devised by J.A.Kitchener and S. Ignatowicz (Ref 10, Fig 1) was therefore employed, yet not the quantity of the volatile product but the hydrogen content of the gas mixture in equilibrium was determined, i.e. by measuring the electromotive force between two hydrogen electrodes, one being saturated with pure hydrogen and the other with a mixture of hydrogen and argon. In order to check the precision of the apparatus, the authors measured the temperature dependence of the equilibrium constant of zinc-oxide reduction (Fig 2, I). Herefrom Card 1/2

The Equilibrium Between Zinc Tungstate and Hydrogen and the Thermodynamic Characteristics of ZnWO

66183 SOV/20-128-5-37/67

it resulted that it was in good agreement with the values obtained by Kitchener and Ignatowicz. For the reaction  $ZnWO_4 + 4H_2 = Zn_{gas} + W + 4H_2O$ , the authors calculated the reaction constant  $K_p$  and determined the phase composition of the reaction products with the help of radiography. Experimental results are listed in table 1, and figure 2, II indicates the dependence of  $lgK_p$  on 1/T.  $\Delta H_{298}^O = -327.0$  kcal/mol,  $\Delta S_{298}^O = -110.66$  cal/mol.degree,  $\Delta Z_{298}^O = -285.1$  kcal/mol were computed by a method developed by M. I. Temkin and L. A. Shvartsman (Ref 16). There are 2 figures, 1 table, and 16 references, 10 of which are Soviet.

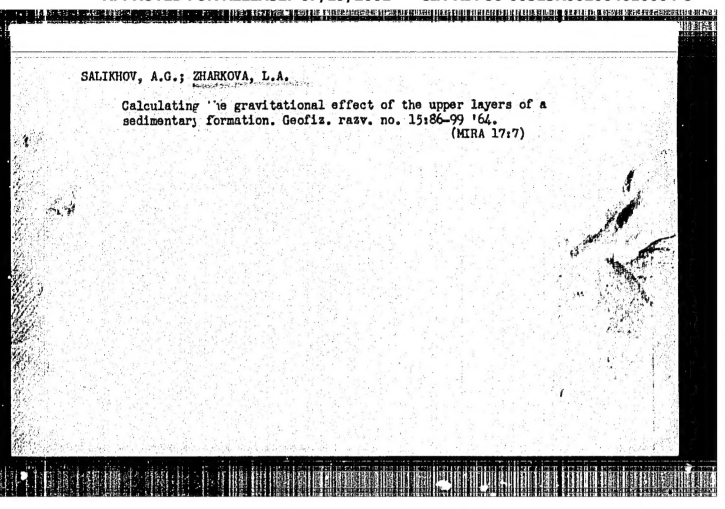
ASSOCIATION:

Moskovskiy gosudarstvennyy universitet im. M. V. Lomonosova (Moscow State University imeni M. V. Lomonosov)

SUBMITTED:

July 2, 1959

Card 2/2



5(4), 24(8)

AUTHORS: Zharkova, L. A., Rezukhina, T. N.

A CONTRACTOR OF THE PARTY OF TH

507/76-32-10-1/39

TITLE:

The Specific Heat of the Nickel, Strontium and Zinc Tungstates and the Barium and Strontium Molybdates at High Temperatures (Teployemkost' vol'framatov nikelya, strontsiya i tsinka i molibdatov bariya i strontsiya pri vysokikh temperaturakh)

PERIODICAL:

Zhurnal fizicheskoy khimii, 1958, Vol 32, Nr 10, pp 2233-2235 (USSR)

ABSTRACT:

The determinations mentioned in the title were carried out in the molar calorimeter within the temperature ranges of 683,2-293,2°K to 1125,2-293,2°K. The scheme, the method employed as well as other details were already described (Ref 1). Data on the technique of preparation and analysis are given. The results obtained are given in a table. In the table the mean values of the specific heat are given for each temperature range, and so are the comparative values of parallel experiments. Equations for the calculation of the mean specific heat as well as the data obtained using them are mentioned. The mean specific heat  $(\overline{C}_p)$  of all investigated salts varies linearly with the tempera-

Card 1/2

ture within the ranges investigated. The specific heat C was

SOV/76-32-10-1/39

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The Specific Heat of the Nickel, Strontium and Zinc Tungstates and the Barium and Strontium Molybdates at High Temperatures

calculated from the mean specific heat according to the equation

 $c_{p} = \frac{d\left[\overline{c}_{p} \left(T - 293, 2\right)\right]}{dT}$ 

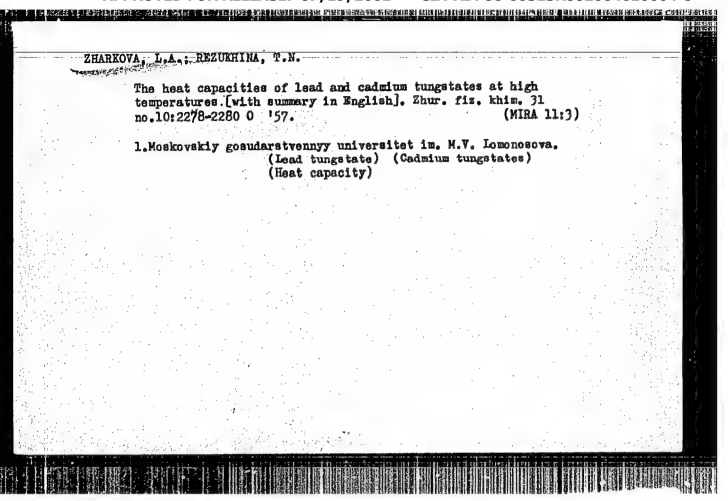
The function  $C_p(T)$  is given individually for the chemical compounds investigated. The authors thank Professor S. M. Skuratov for his advice. There are 1 table and 2 references, 2 of which are Soviet.

ASSOCIATION: Moskovskiy gosudarstvennyy universitet im. M. V. Lomonosova

(Moscow State University imeni M. V. Lomonosov)

SUBMITTED: January 30, 1957

Card 2/2



1. 1 14 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	THE STATE OF THE S
ZHARKOUA, L	, A . 76-10-15/34
AUTHORS:	Zharkova, L.A., Rezukhina, T.N.  Specific Heat of Lead and Cadmium Tungstate at High Temperatures  (Teployemkosti vol'framatov svintsa i kadmiya pri vysokikh temperaturakh)
PERIODICAL:	Zhurnal Fizicheskoy Khimii, 1957, Vol. 31, Nr 10, pp. 2278-2280  Zhurnal Fizicheskoy Khimii, 1957, Vol. 31, Nr 10, pp. 2278-2280  Data for the specific heat within the range of from 800 - 20°C  Data for the specific heat was determined according to the are given here. The specific heat was determined according to the method for mixing in a massive calorimeter. The description of method for mixing in a massive calorimeter. The description of the device is found in M.M. Popov's "Termometriya i kalorimetrithe device is found in M.M
Card 1/2	The mean specific near $p$ range amounts to: $\overline{C}_{p}, PbWO_{4} = 0.06566 + 1.034 \cdot 10^{-5} \text{ T (accuracy } \pm 0.03 \%)$ $\overline{C}_{p}, CdWO_{4} = 0.07754 + 1.9041 \cdot 10^{-5} \text{ T (accuracy } \pm 0.10 \%)$

APPROVED FOR RELEASE: 07/19/2001 CIA-RDP86-00513R002064610004-8"

Specific Heat of Lead and Cadmium Tungstate at High Temperatures 76-10-15/34

The real specific heat can be expressed by following equations:

= 0,06263 + 2,068 . 10<sup>-5</sup> T

<sup>C</sup>p, CdWO<sub>A</sub> = 0,07195 + 3,8082 -10<sup>-5</sup> T

T - is the absolute temperature. There are 1 table and 3 Slavic

ASSOCIATION:

Moscow State University imeni M.V. Lomonosov

(Moskovskiy gosudarstvennyy universitet imeni M.V. Lomonosova)

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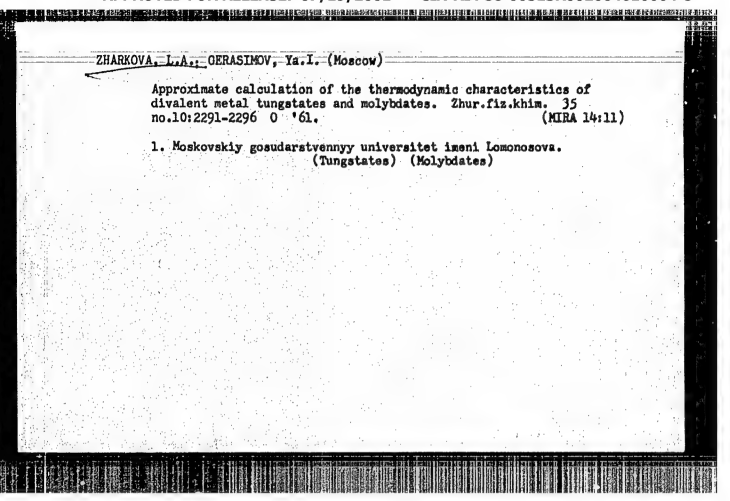
July 18, 1956

AVAILABLE:

Library of Congress

Card 2/2

ZHARKOVA, L. A., Cand Chem Sci (diss) -- "Thermodynamic investigation of certain wolframates and molybdates". Moscow, 1960. 13 pp (Moscow State U im M. V. Lomonosov, Chem Faculty, Chair of Phys Chem), 120 copies (KL, No 14, 1960, 127)

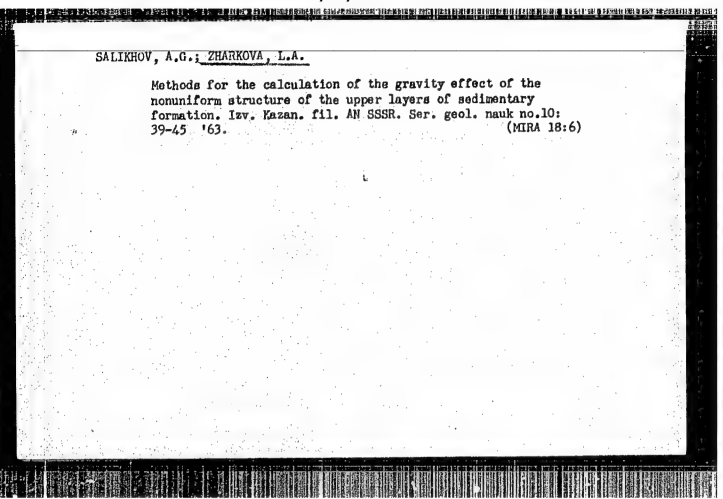


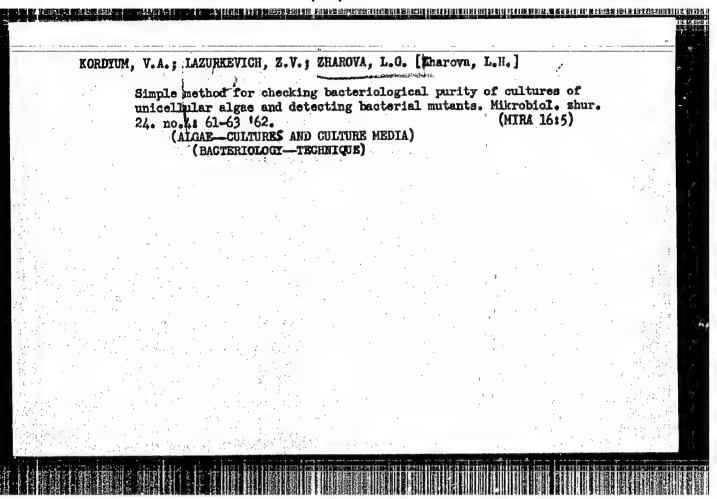
SALIKHOV, A.G.; ZHARKOVA, L.A.; KUZNETSOV, G.Ye.

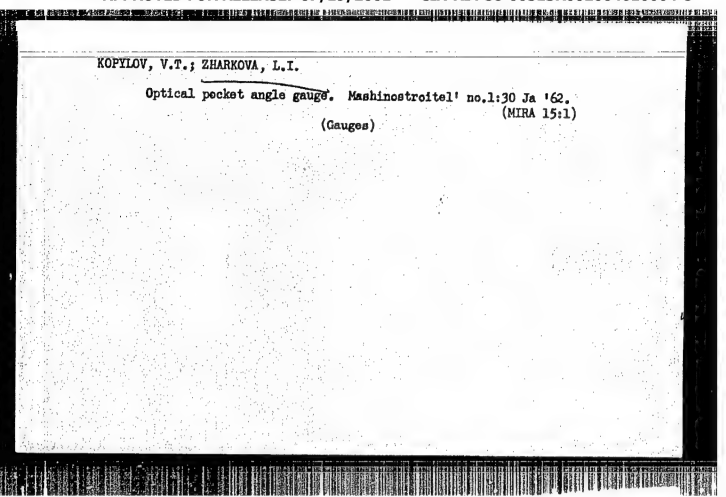
Fast method for the determination of the elements of occurrence and the gravity effect of disturbing masses having the form of a bench. Izv. Kazan. fil. AN SSSR. Ser. geol. nauk no.10-18-26 163.

Interpretation of the materials of detailed gravimetric surveying in Tatarstan. Ibid.: 144-150

(MIRA 18:6)







B/183/61/000/005/002/003 B101/B110

15 5560

Zharkova, M. A., Rassolova, E. A., Kudryavtsev, G. I.,

**李明氏是因为医生的性性**。例如我们的对我们的对称,但这种思维的理解,我们是这是否则可以是可以且们对此的。

AUTHORS Klimenkov, V. S.

Copolymerization of acrylonitrile and 2-methy1-5-vinyl TITLE:

pyridine in aqueous sodium thiocyanate solution

Khimicheskiye volokna, no. 5, 1961, 13 - 17 PERIODICAL:

The authors attempted to improve the quality of acrylonitrile fibers by means of pyridine derivatives. Previous papers (Khim. volckna, no. 3, 15 (1960); ibid., no. 6, 15 (1960)) dealt with the copolymerization of acrylonitrile (AN) and  $\alpha$ -vinyl pyridine ( $\alpha$ -VP). In the present paper, the system AN - 2-methyl-5-vinyl pyridine (MVP) was studied, since a simple method of producing MVP has been developed in the Soviet Union. 50% sodium thiocyanate proved to be an optimum solution for copolymerization. Experiments at room temperature and 70°C showed that the formation of sufficiently concentrated homogeneous spinning solutions (12 - 15%) with a maximum ratio AN:MVP = 85:15 is limited due to the poor solubility of MVP. Copolymerization of AN and MVP is analogous to that of AN and

Card 1/4

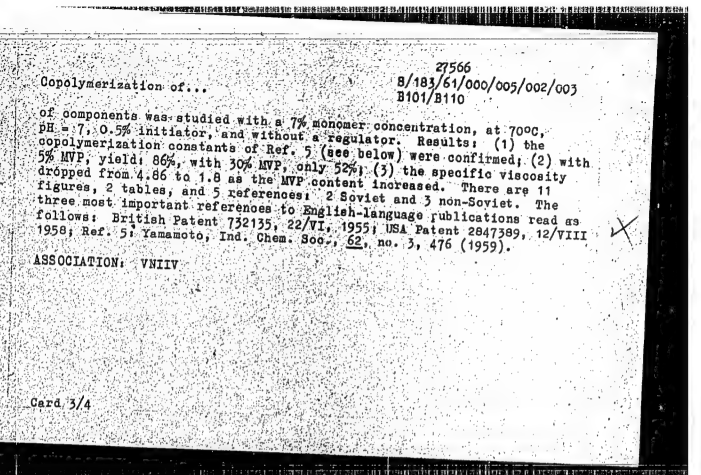
APPROVED FOR RELEASE: 07/19/2001 CIA-RDP86-00513R002064610004-8"

27566 8/183/61/000/005/002/003 B101/B110

Copolymerization of ...

The yield after 60 min is 60 - 65%. The reaction rate drops linearly with the time of polymerization. Fig. 4 shows that the pH of the medium exerts a considerable effect upon the yield. These data are not in agreement with those cotained by Yamamoto (see below). Only in acid media does the specific viscosity depend on pH; in alkaline media it is constant. The initiator used in copolymerization was azodinobutyric acid dinitrile. The polymerization rate was found to be a linear function of the square root of the initiator concentration. With 0.05% initiator (optimum concentration), the polymer yield after 1.5 hr is 75 - 80%. A rise in temperature (from 60 to 80°C) accelerates the process. 70°C is optimum for a 7% monomer solution, since the polymerization rate is not high enough as to cause overheating. The activation energy is 14.5 kcal/mole. To obtain optimum spinning solutions, the specific viscosity should not exceed 1.0 - 1.2. Therefore, experiments were made with various regulators: monoethanol amine, thiourea, thymol, lauryl mercaptan, diproxide (= dipropyl xanthogenatedisulfide), thiourea dioxide. Monoethanol amine was the only substance to affect the molecular weight of the polymer. 0.7% of monoethanol amine (with α-VP only 0.2%) was required to obtain AN-MVP copolymers of the desired viscosity. The effect of the ratio

Card 2/4



ZHARKOVA, L.P.; MOVSHOVICH, T.Kh.; T.E. VA, L.G.; ROZITIS, T.Ya.;

GOLUBISOV, I.Ys., otv. red.: E.GACHEVA, G.V., red.;

ROMANOVA, S.P., tekhn. red.

[Rural K-40/80 crosstar automatic telephone exchanges]

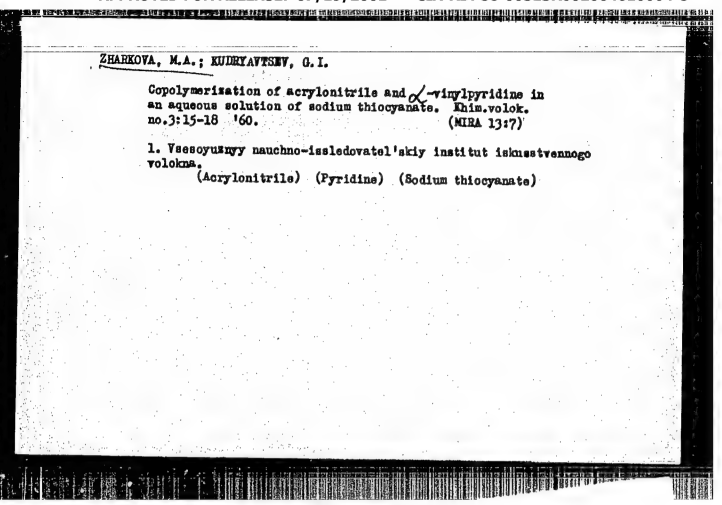
Sel'skie koordinathye ATC K-40/80; informatsionnyi sbornik,

Moskva, Svias'12dat, 1963. 109 p. (NIRI 16:10)

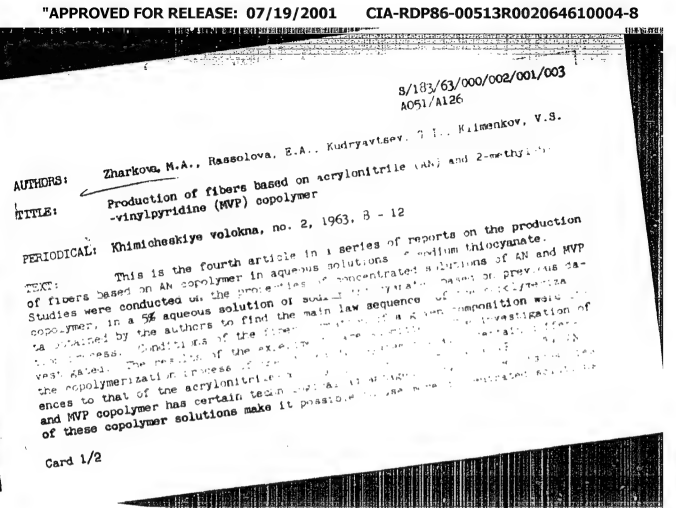
1. Nauchno-iseledovatel'skiy institut gorodskoy i sel'skoy
telefonnoy svyazi Ministerstva svyazi SSSR (for Zharkova,

Movshovich, Frolova). 2. Gosudarstvonnaya elektrotekhnicheskaya fabrika, Riga (for Rudzitis).

(Telephone)



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CIA-RDP86-00513R002064610004-8" APPROVED FOR RELEASE: 07/19/2001

Production of fibers based on acrylonitrile

S/183/63/000/c02/col/col

or polymers of a higher molecular weight. The AN-MVP system allows for a wider range of the polymer concentration change that the Ah-a-Vr system allows for a wider with a specific viscosity above 2. Experiments showed the optimum specific viscosity to be 1.25 - 1.5. A slight temperature elevation of the solution reduces the latter. Investigated solutions of 0.8, 1.25, 1.48 intrial specific viscosity left to stand, did not selatinize at 25 0.70 C, even when specific viscosites, remained constant. There are 5 figures and 1 table.

ASSOCIATION: VNIIV

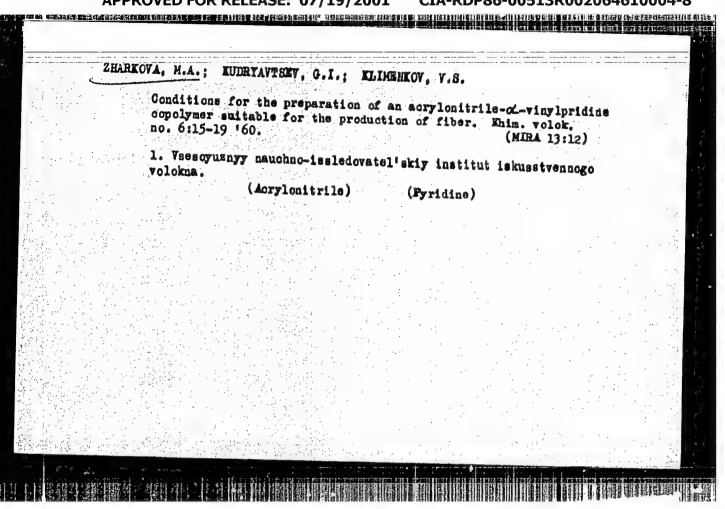
Card 2/2

Card 2/2

BUNAREVA, 2.3.; DIURNBAUM, V.S.; DOROKHINA, I.S.; ZHARKOVA, M.A.; KLIMENKOV, V.S.

Fibers based on mixtures of adrylonitrile polymers. Khim.volok no.6;10(MIRA 17;1)

1. Vsesoyusnyy hauenno-issledovatel skiy institut iskusstvennogo volokna.



ZHARKOVA

5/183/60/000/03/06/007 B020/B054 8206L

15.5560

The state of

Kudryavtsev, G. I.

Copolymerization of Acrylonitrile and  $\alpha$ -Vinyl Pyridine in AUTHORS:

Aqueous Sodium Thiocyanate Solution TITLE:

Khimicheskiye volokna, 1960, No. 3, pp. 15-18 PERIODICAL:

TEXT: As no publication data are available on the copolymerization of acrylonitrile (AN) with  $\alpha$ -vinyl pyridine ( $\alpha$ -VP) in aqueous sodium thiooyanate solutions, the present paper studies the principal rules governing the process of producing a thread-forming copolymer with low vinyl pyridine content. Table 1 shows the change in composition of the copolymer from the initial ratio of monomers in copolymerization; it was found that, in agreement with theoretical calculations, the copolymer obtained always exhibits an increased a-VP content. Fig. 1 shows the dependence of the copolymer yield on the initial concentration of monomers in the solution, Fig. 2 the dependence of the initial rate of polymerization on the monomer concentration in the solution, Fig. 3 the dependence of the monomer consumption on time at different concentrations of the initiator

Card 1/2

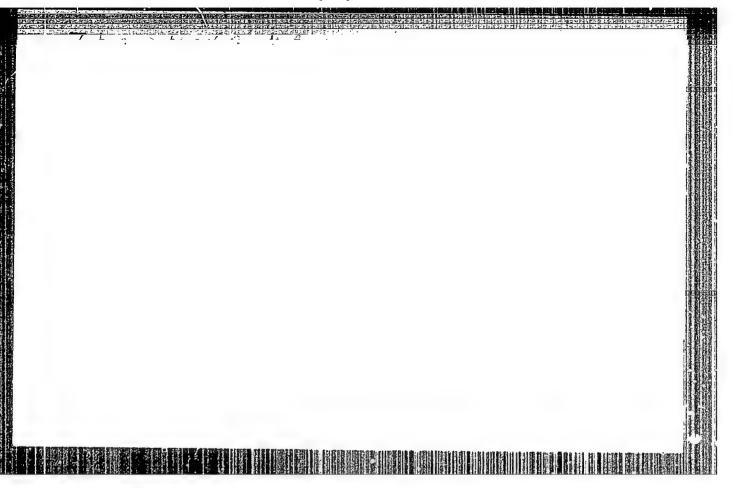
的复数形式不好,这种的现在分词,这种种的一种的一种的一种,但是一种的一种,是一种种的一种,这种的一种,我们可以是一种的一种的一种的一种的一种,我们可以使用的一种

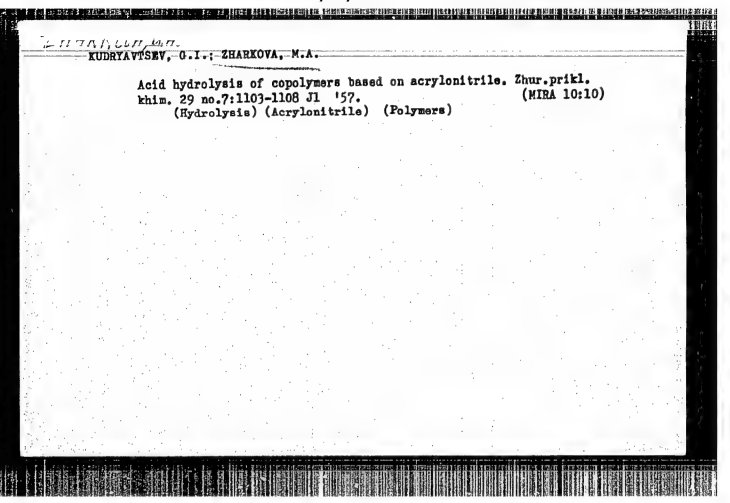
Copolymerization of Acrylonitrile and a-Vinyl S/183/60/000/03/06/007 Pyridine in Aqueous Sodium Thiocyanate Solution B020/B054

at a ratio AN:  $\alpha$ -VP = 95: 5% by weight, and Fig. 4 the same dependence on the polymerization rate of AN with  $\alpha$ -VP is indicated in Table 2. Table 3 shows the influence of regulators (lauryl mercaptan, thiourea, influence of the monoethanol amine amount on the copolymer yield. It is shown that the reaction rate is proportional to the initial concentration of the monomer mixture and the square root of the initial used, the methods of investigation, and the determination of copolymer composition. E. A. Rassolova cooperated in working out the methods. British, and 2 French.

ASSOCIATION: VNIIV (All-Union Scientific Research Institute of Fibers)

Card 2/2





ACC NR: AP7000329 (A) SOURCE CODE: UR/0413/66/000/022/0077/0077

INVENTOR: Kudryavtsev, G. I.; Zharkova, M. A.; Romanova, T. A.; Klimenkov, V. S.

ORG: none

TITLE: Method of preparing modified polyacrylonitrile fiber. [announced by the All-Union Scientific Research Institute of Synthetic Fiber (Vsesoyuznyy nauchnoissledovatel'skiy institut iskustvennogo volokna)] Class 29, No. 188617

SOURCE: Izobreteniya, promyshlennyye obraztsy, tovarnyye znaki, no. 22, 1966,

TOPIC TAGS: polyacrylonitrile, hydrazine, synthetic material

ABSTRACT: A method of preparing modified polyacrylonitrile fiber is introduced. To raise the chemical and thermal resistance of the fiber, it is treated in a hydrazine solution and heat treated in an inert-gas medium at 150—200C. [Translation]
SUB CODE: 11/SUBM DATE: 17Sep64/

Card 1/1

UDC: 677. 494. 745. 32:546. 171. 5

8747B \$/183/60/000/006/003/005 B020/B058

AUTHORS:

Zharkova, M. A., Kudryavtsev, G. I., Klimenkov, V. S.

TITLE:

Study of the Conditions of Copolymer Production From Acrylonitrile With Alpha Vinyl Pyridine, Suitable for Fibration

PERIODICAL:

Khimicheskiye volokna, 1960, No. 6, pp. 15-19

TEXT: The paper reports on the results of studies concerning: a) copolymerization of acrylonitrile (AN) with  $\alpha$ -vinyl pyridine ( $\alpha$ -VP) for the purpose of producing a copolymer with predetermined molecular weight and the determination of the optimum concentration of the spinning solution, b) the determination of the optimum concentration of the salt solution, c) the conditions for the production of suitable spinning solutions, and d) the trial formation in precipitating baths with aqueous salt solutions and the study of the physical and mechanical properties of the fiber obtained. In copolymerization, the molecular weight of the copolymer is influenced by the amount of the initiator (azo-dissobutyric acid-dinitrile), the temperature, type of solvent and amount of the regulator (monosthanol amine). Copolymers with a ratio AN:  $\alpha$ -VP of 85: 15 and 90: 10 weight% Card 1/3

87478

Study of the Conditions of Copolymer Production 5/183/60/000/006/003/005 From Acrylonitrile With Alpha Vinyl Pyridine, B020/B058 Suitable for Fibration

were studied. The influence of the amount of regulator on the change in time of the intrinsic viscosity (Fig. 1), and the dependence of the intrinsic viscosity on the regulator concentration (Fig. 2) are determined. The change of the intrinsic viscosity of the solution in dependence on the amount of initiator used is mentioned in Figs. 3 and 4. It can be seen from Fig. 5 that with rising temperature, the intrinsic viscosity of the co-polymer produced drops from 2.5 at 60°C to 1.3 at 75°C. The dependence of the intrinsic viscosity of the copolymer on the initial concentration of the monomer mixture (Fig. 6) shows that the probability of a chain rupture through the solvent increases with sinking concentration of the monomers in the solution. As may be seen from the tabulated data concerning the conditions of the copolymerization of AN with  $\alpha$ -VP in the production of spinning solutions, the rate of polymerization in 45 to 50% sodium thiocyanate, under otherwise equal conditions, is always tha same and the copolymers have the same intrinsic viscosity (1.39 to 1.4). Fig. 7 shows the dependence of the viscosity of a concentrated sodium thiocyanate solution on the intrinsic viscosity of the copolymer. It can be seen from Fig. 8 that at an intrinsic viscosity of 1.38, 10.5% to 11.2% Card 2/3

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87478

Study of the Conditions of Copolymer Production S/183/60/000/006/003/005 From Acrylonitrile With Alpha Vinyl Pyridine, B020/B058

solutions are suitable for the shaping of the fiber, and at an intrinsic viscosity of 0.97, 15% solutions. The fiber produced under the optimum conditions determined had the following values: metric number 3970, breaking length 25.6 km, elongation 32%; the fiber can be dyed well with a cetate and alkaline dyes. There are 8 figures, 1 table, and

ASSOCIATION: VNIIV (All-Union Scientific Research Institute of Synthetic Fibers)

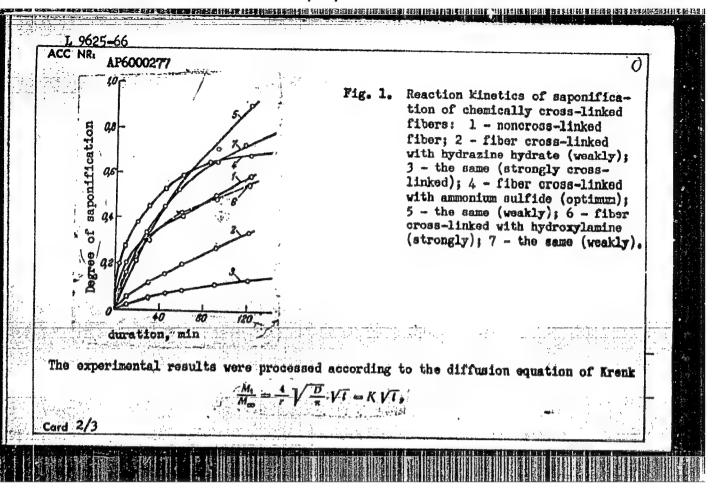
Card 3/3

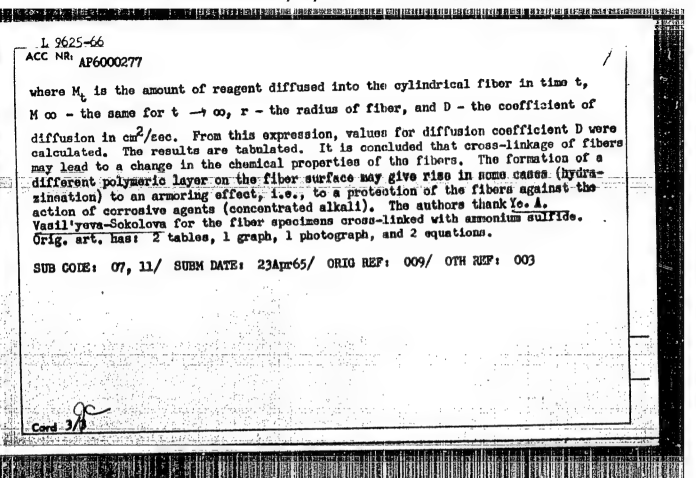
ZHARKOVA, M.A.; RASSOLOVA, E.A.; KUDRYAVTSEV, G.I.; KLIMENKOV, V.S.

Copolymerization of acrylonitrile and 2-methyl-5-vinylpyridine in an aqueous solution of sodium thiocyanate. Khim.volok. no.5:12-17 '61. (MIRA 14:10)

1. Vegeoyuznyy nauchno-issledovatel'skiy institut iskusstvennogo volokna. (Acrylonitrile) (Pyridine) (Polymerization)

ACC NRI AP6000277	SOURCE CODE: UR/0183/65/000/005/0013/0015
AUTHORS: Kudryavtsev, G.	I.; Romanova, T. A.; Zharkova, H. A.; Klimenkov, V. S.
ORG: WHIV W	413
TITLE: Some chemical pro	operties of cross-linked PAN (polyacrylonitrile) fibers
SOURCE: Khimicheskiye vo	olokna, no. 5, 1965, 13-15
TOPIC TAGS: fiber, acrylplastic, synthetic fiber	ionitrile, acrylonitrile polymer, acrylic resin, polymer,
saponification of chemica The study was undertaken	to extend the presently available literature data on the coss-linked PAN-fibers (polyacrylonitrile fibers) (1994) to extend the presently available literature data on the coss-linked PAN-fiber, as compiled by G. I. Kudryavtsev, T.
A. Matyash, M. A. Zharkov The saponification kinet Ifiable groups in PAN-fil	va, and V. S. Klimenkov (Khi., volokna, No. 4, 13, 1961).  los at 1000 of nitrile and other nitrogen-containing sapon- per cross-linked by hydrazine hydrate, hydroxylamine, and
uring the amount of amnor	lied. The degree of saponification was determined by meas- nia released by the fibers after treatment with 40% NaVH
	tal results are presented in tables and graphs (see Fig. 1).
ard 1/3	UDC: 677.494.745.32:061.3



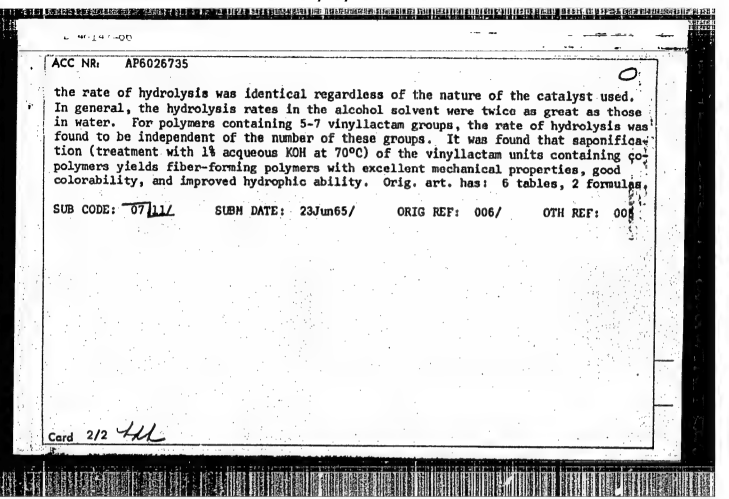


ZHARKOVA, M.A., ROMANOVA, T.A.

Chemical-resistant fibers with a base of acrylonitrile. Khim. volok. no.5:77 165. (MIRA 18:10)

1. Vsesoyuznyy nauchno-issledovatel skiy institut iskusstvennogo volokna.

L 46147-66 EWT(m)/EWP(1)/T LJP(c) WW/RM VSOURCE CODE: UR/0183/66/000/003/0012/0015	7
ACC NR: AP6026735 (A)  AUTHOR: Kudryavtsev. G. I.; Rassolova, E. A.; Romanova, T. A.; Zharkova, H. A.; Postil'yeva-Sokolova, Ye. A.	
ORG: VNIIV  TITLE: Preparation and modification of fiber-forming polymers made of vinyllactemunits containing acrylonitrile	
SOURCE: Khimicheskiye volokna, no. 3, 1966, 12-15  TOPIC TAGS: polyacrylonitrile, synthetic fiber, copolymerization, catalytic polymerization, polymerization kinetics, copolymer  ABSTRACT: The kinetics of the hydrolysis of polyvinylcaprolactam and acrylonitrile  ABSTRACT: The kinetics of the hydrolysis of polyvinylcaprolactam and acrylonitrile  vinylcaprolactam copolymer was studied. The object of the work was to prepare read  vinylcaprolactam copolymer was studied. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and hydrophylic fibers. The hydrolysis constants were measured at 100  ly colorable and	it- nic



# "APPROVED FOR RELEASE: 07/19/2001

(MIRA 18:5)

YELISEYEVA, V.I.; ZHARKOVA, N.G.; CHUBAROVA, A.V.; ZUEOV, P.I. Emulsion polymer:zation of lower alkyl acrylates. Vysokom.soed. 7 no.1:156-162 Ja '65. (MIRA 18

1. Institut fizicheskoy khimii AN SSSR.

CIA-RDP86-00513R002064610004-8" APPROVED FOR RELEASE: 07/19/2001

L 13291-66 EWT(m)/EWP(1) RM ACC NR: AP6000324 SOURCE CODE: UR/0286/65/000/021/0011/0011 INVENTOR: Zharkova, N. I.; Zamarayev, A. P.; Koroleva, Ye. S. ORG: none TITLE: A method for preparation of a catalyst to produce vinyl benzene No. 175927 SOURCE: Byulleten izobreteniy i tovarnykh znakov, no. 21, 1965, 11 TOPIC TAGS: vinyl plastic, polymerization catalyst, aromatic hydrocarbon ABSTRACT: This Author's Certificate introduces a method for preparing a catalyst to produce viny1 benzene. Diethyl benzene is dehydrated by mixing and preforming the active components. The product yield is increased and a stable catalyst is produced by preparing it from two layers with the following composition: upper layer-68.3 %, 15 % magnesium oxide, 4.4 % copper oxide, 12.3 % sodium carbonate, lower layer--72.7 % iron oxide, 16 % magnesium oxide, 4.8 % copper oxide, 6.6 % potassium carbonate. SUB CODE: 07/ SUBM DATE: 30Nov62/ ORIG REF: 000/ OTH REF: 000 Card 1/1 66.097.3 : 547.538.1.07

KIEYNOVSKAYA, M.A.; SOBOLEVSKIY, M.V.; ZHARKOVA, N.M.

Composition and properties of liquid polyorganosiloxanes as dependent on the method of synthesis employed. Report No.1: Composition and properties of polymethylphenylsiloxanes obtained by cohydrolysis method. Plast.massy no.7:27-31 '62. (MIRA 15:7)

(Silicon organic compounds)

38719 \$/191/62/000/007/006/011 B124/B144

AUTHORS:

Kleynovskaya, M. A., Sobolevskiy, H. V., Zharkova, N. E.

多类性多种的表现的。一种,这种是一种,我们是一个人,我们是一个人,我们是一个人,我们是一个人,我们是一个人,我们是一个人,我们是一个人,我们是一个人,我们是一个人,我们就是一个人

TITLE:

Investigation of composition and properties of liquid polyorganosiloxanes as depending on the method of synthesis. Communication I. Investigation of composition and properties of polymethyl phenyl siloxanes produced by cohydrolysis

PERIODICAL: Plasticheskiye massy, no. 7, 1962, 27-31

TEXT: The authors made the first attempt to determine the composition of polyorganosiloxanes resulting from various methods of synthesis, by means of combined molecular and vacuum rectification in spray and packed towers. An apparatus developed by F. W. Melpolder et al. was used. Separation is conducted under high vacuum. The apparatus may work either intermittently or continuously. With infinite reflux the efficiency of the stills is 0.75. The composition of polymethyl phenyl siloxanes got by cohydrolysis of a methyl phenyl dichlorosilane-trichlorosilane mixture (3:2.2) at 96-100°C in acid solution is investigated. The product was thermally

Card 1/3

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Investigation of composition and ...

stabilized in nitrogen flow, distilled at 0.1-0.3 mm Hg, and collected in four fractions. Apart from the distillation residue disregarded, the reaction product is mainly a mixture of linear polymethyl phenyl siloxanes having the composition (CH<sub>3</sub>)<sub>3</sub>Si[OSiCH<sub>3</sub>C<sub>6</sub>H<sub>5</sub>]<sub>n</sub>OSi(CH<sub>3</sub>)<sub>3</sub> (n = 1-5). Fraction I is a mixture of low-boiling linear polymethyl phenyl siloxanes with 3-4 Si atoms per molecule, fraction II consists mainly of linear methyl phenyl tetrasiloxane with small amounts of methyl phenyl tri- and methyl phenyl pentasiloxane, fraction III of linear methyl phenyl siloxane with 5 Si atoms per molecule, and fraction IV of linear methyl phenyl siloxunes mixed with 6 and 7 Si atoms per molecule besides small amounts (3-9%) of cyclic methyl phenyl siloxanes. Four linear polymethyl phenyl siloxanes were isolated and characterized, the first three of which have not previously been described in publications: 1,1,1,3,5,7,7,7-octamethyl-3,5-diphenyl tetrasiloxane; 1,1,1,3,5,7,9,9,9 nonamethyl-3,5,7-triphenyl pentasiloxane; 1,1,1,3,5,7,9,11,11,11-decamethy1-3,5,7,9-tetrapheny1 hexasiloxane, and 1,1,1,3,5,5,5-heptamethy1-3phenyl trisiloxane. There are 1 figure and 3 tables.

Card 2/3

Investigation of composition and ...

S/191/62/000/007/006/011 B124/B144

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The most important English-language references are: F. W. Melpolder et al., Anal. Chem. 27, No.6, 974 (1955); B. Okawara et al., Bull. Chem. Soc. Japan 30, 608 (1957); H. I. Waterman et al., J. Appl. Chem. 8, No. 10, 625 (1958).

Card 3/3

10204 \$/191/62/000/009/005/012 B101/B144

AUTHORS:

Kleynovskaya, M. A., Sobolevskiy, M. V., Krasovskaya, T.

Zharkova, N. Ma

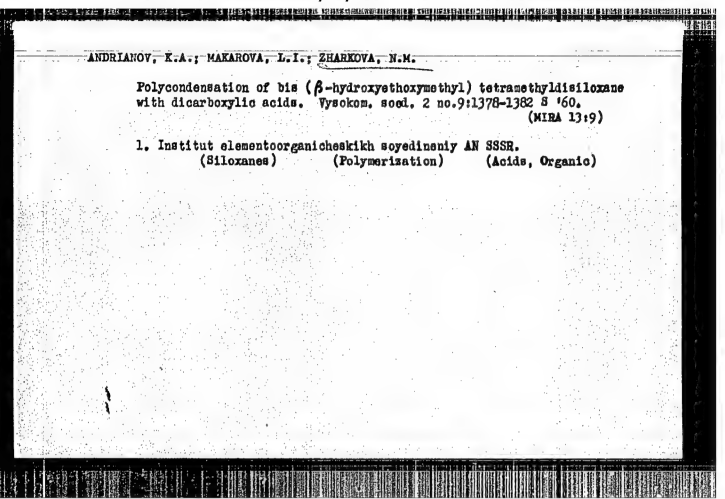
Dependence of the composition and properties of liquid TITLE

polyorganosiloxanes on their mode of production

Plasticheskiye massy, no. 9, 1962, 19 - 24 PERIODICAL:

TEXT: The composition and properties of polymethyl-phenyl siloxenes got by cohydrolysis and subsequent catalytic regrouping in the presence of Kil clay were studied as follows: Aqueous solutions of methyl-phenyl dichlorosilane, dimethyl dichlorosilane and trimethyl chlorosilane in the molar ratio 3:1:2.2 were cohydrolyzed at 60-65 C. The cyclic byproducts developed were regrouped with 8% Kil clay as catalyst at 50°C (6 hr) into linear compounds. The reaction product was fractionated and investigated. Predominantly linear polymers having the general formula: (CH<sub>3</sub>)<sub>3</sub>Si[OSiCH<sub>3</sub>C6<sup>H</sup>5]<sub>n</sub>[OSi(CH<sub>3</sub>)<sub>2</sub>]<sub>m</sub>OSi(CH<sub>3</sub>)<sub>3</sub> resulted. In the products distilled within the limits of 380°C/0.1-0.5 mm Hg, n was 0,1,...6; m was m was 0,1;...7. The content of cyclic compounds did not exceed 0,1,2; n +Card 1/2

era estructus el la cultura de la compania de la c 5/191/62/000/009/005/012 Dependence of the composition. B101/B144 4%. The product contained 1.2% hexamethyl disiloxane, 1.8% & w-hexamethylpolydimethyl siloxanes, 26% &, w-hexamethyl-polymethyl-phenyl siloxanes, 45% co, w-hexamethyl-polydimethyl-polymethyl-phenyl siloxanes. The 23% of nondistillable residues seem to be composed of high-boiling polymers of the latter type. For 11 compounds of the given general formula, 10 of which were synthetized for the first time, b.p. (C/mm Hg), softening point (C),  $n_D^{20}$ ,  $d_4^{20}$  and  $\eta_{20}$  (centistokes) are given respectively as follows: n=1, m = 0:78-79/0.5, -, 1.4470, 0.9118, 2.55; n = m = 1:87-88/0.5, -75, 1.4393, 0.9244, 3,75; n = 1, m = 2:105/0.5, -95, 1.4363, 0.9355, 3.85; n = 2, m = 0; 130-132/0.5, -75, 1.4775, 0.9761, 7.05; n = 2, m = 1; 147-149/0.5, -70, 1.4670, 0.9786, 7.77; n = m = 2; 162/1.0, -60, 1.4605, 0.9807, 8.50; n = 3, m = 0; 180/0.5, -60, 1.4950, 1.0132, 15.71; n = 3, 1.0331, 29.17; n = 4, m = 2; -60, 1.4930, 1.0327, 27.55; n = 5, m = 2; -60, 1.4930, 1.0327, 27.55; n = 5, m = 2; -8, regular connection exists between the physicochemical properties and the content of dimethyl- and methyl-phenyl siloxane links. There are 5 Card 2/2



8/190/60/002/009/010/019 alm 2103, 2209 B004/B060 Zharkova, N. M. Makarova, L., I., AUTHORS: Polycondensation of Bis-(β-hydroxy-ethoxy-methyl)-tetra-TITLE: methyl Disiloxane With Dicarboxylic Acids Vysokomolekulyarnyye soyedineniya, 1960, Vol. 2, No. 9, PERIODICAL: pp. 1378-1382 TEXT: The authors studied the condensation of organosilicon diketo carboxylic acids CH3 -SiCH2C6H4CO-C6H4COOH with HOCH2CH2OCH2SiOSICH2OCH2CH2OH at 220°C. Unlike the reaction with ethylene glycol, no cyclic polydimethyl siloxanes were formed. A table shows the Card 1/3

83479

Polycondensation of Bis-(β-hydroxy-sthoxy-methyl)- S/190/60/002/009/00/019 tetramethyl Disiloxane With Dicarboxylic Acids B004/B060

silicen content of the condensates. There occurred neither a cleavage of the siloxane bond in the diketocdicarboxylic acid nor a cleavage of the Si-C bond in organosilicon glycol. As is shown in Fig. 1, the acid number drops during polycondensation while the ester number rises. The polyesters obtained are high-viscous, dark-colored substances well soluble in benzene - alcohol mixture. As is shown by Fig. 2; the viscosity of polyesters rises with the number of dimethyl siloxane groups in diketo dicarboxylic acid. On the reaction of the polyester obtained from diketo dicarboxylic acid (n=5) with hexamethylene diisocyanate, the authors obtained an elastic, rubber-like, cresol-soluble product. The change (increase) in viscosity as dependent on the reaction period is illustrated in Fig. 3. Bis-(β-hydroxy-ethoxy-methyl)-tetramethyl disiloxane also condenses with adipic acid without a cleavage of the Si-C bond to form a polyester. Fig. 4 shows the change in the acid number and ester number during the reaction. There are 4 figures, 1 table, and 3 references: 2 Soviet and 1 US.

Card 2/3

83479

Polycondensation of Bis-(β-hydroxy-ethoxy-methyl)-tetramethyl Disiloxane With

3/190/60/002/009/010/019 B004/B060

Dicarboxylic Acids

ASSOCIATION:

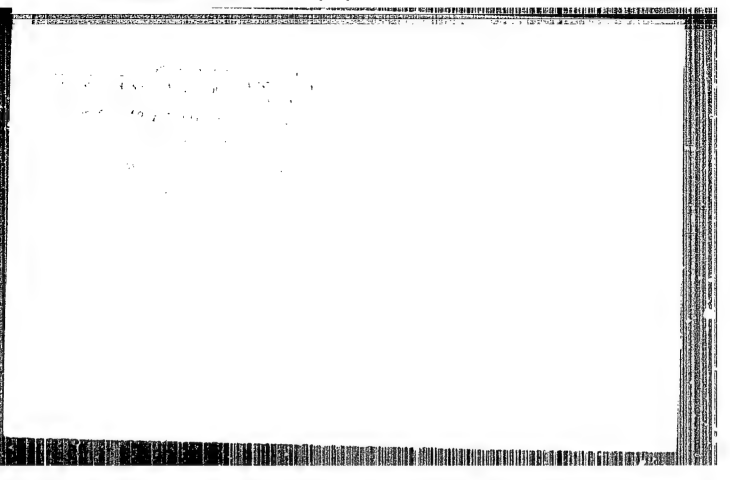
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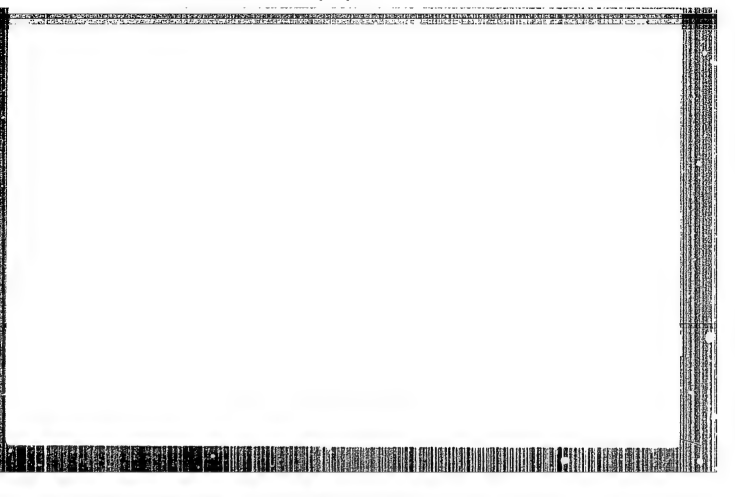
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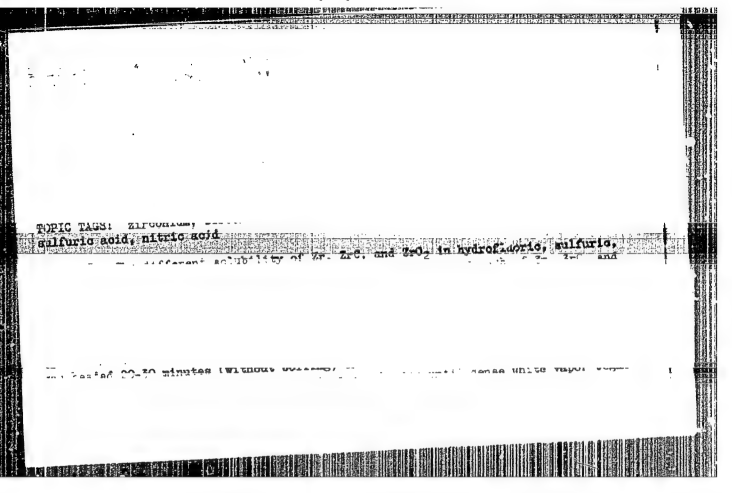
April 11, 1960

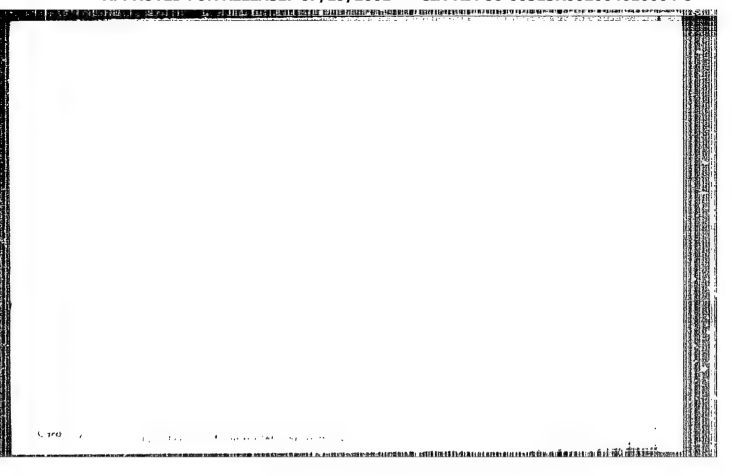
Card 3/3

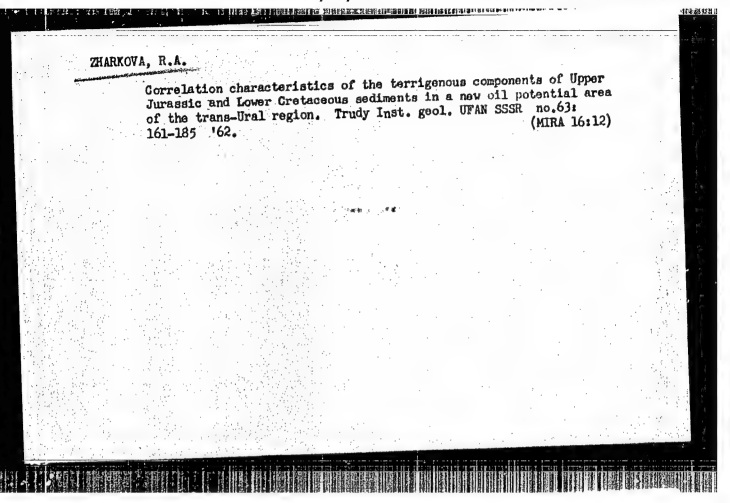
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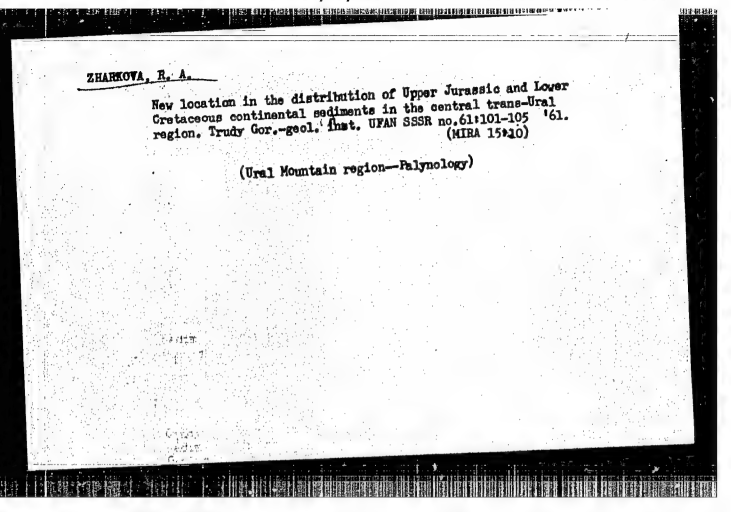


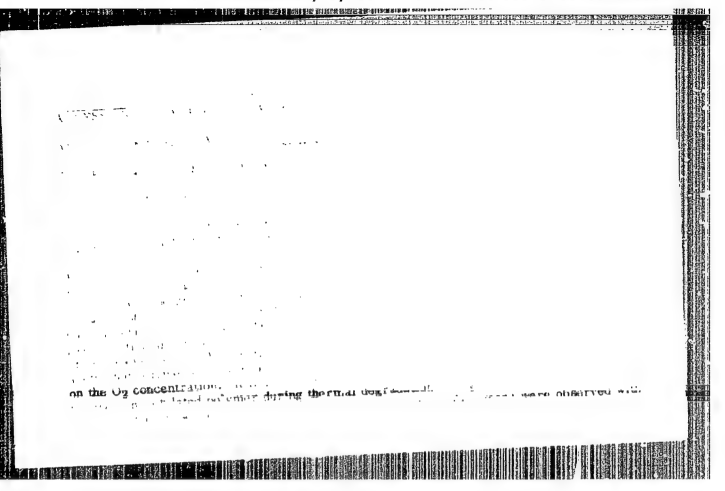


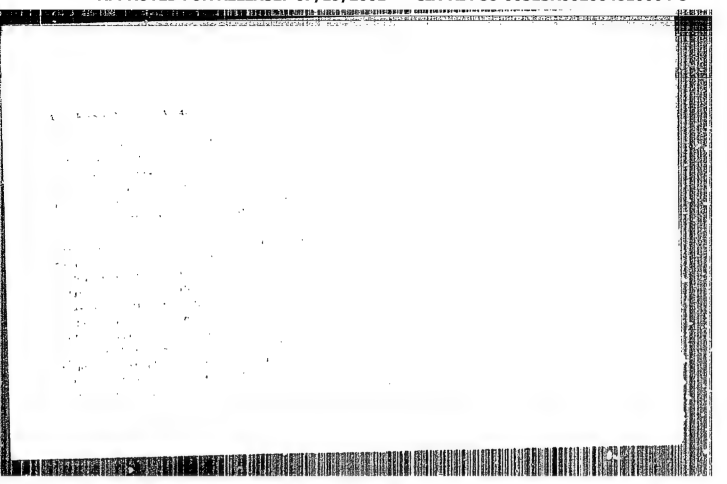


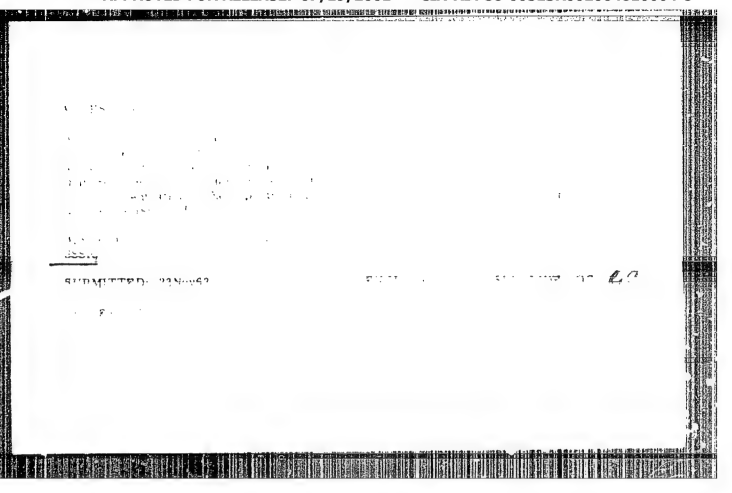


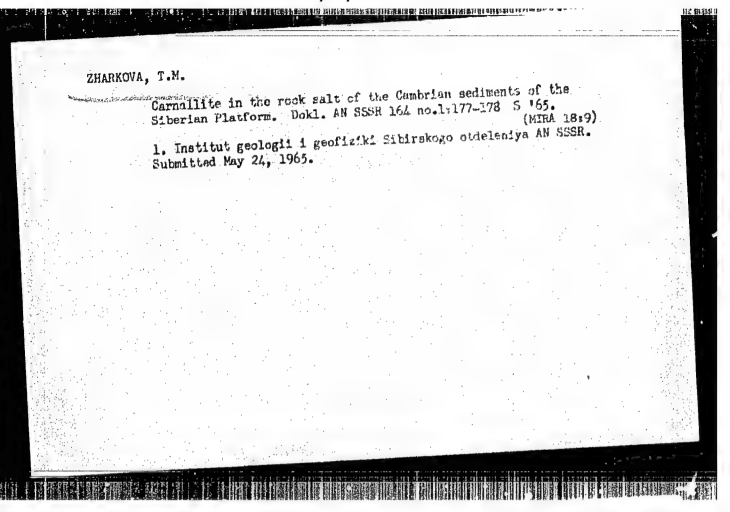


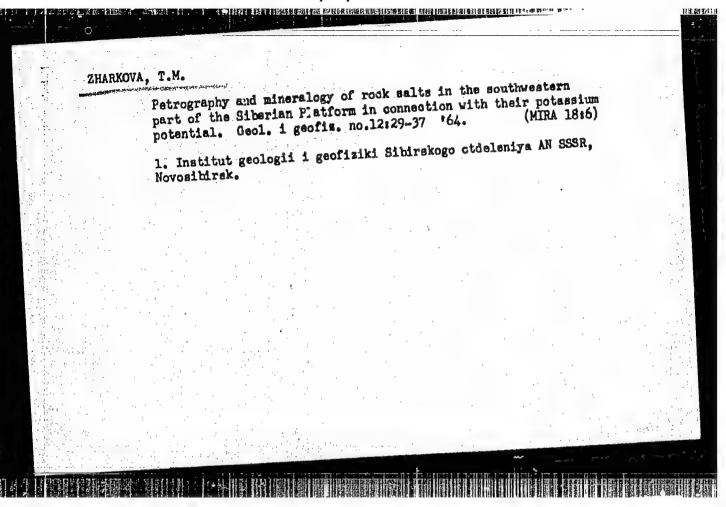


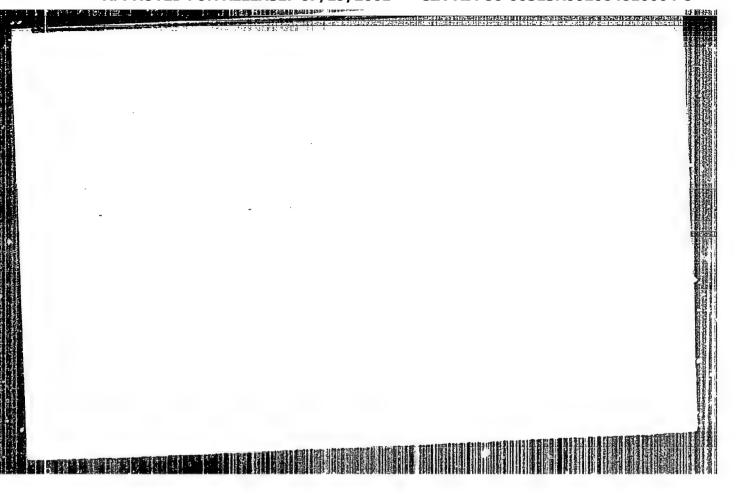


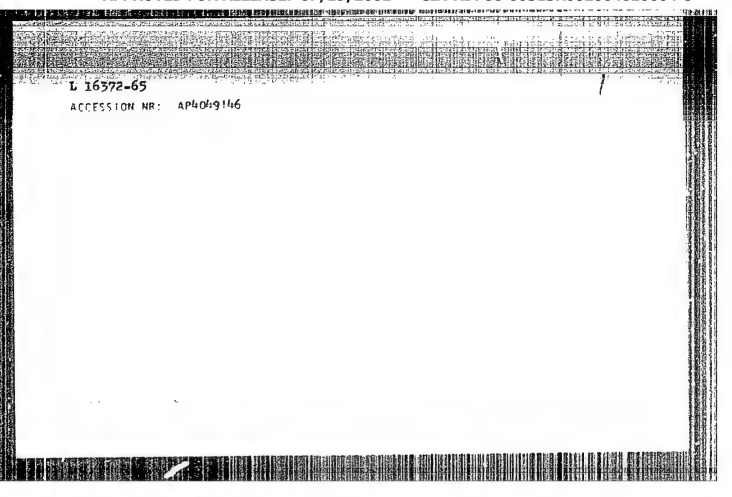












SAMUSEVA, R.G.; ZHARKOVA, R.M.; PLYUSHCHEV, V. Ye.

System Na<sub>2</sub>MoO<sub>4</sub> - Cs<sub>2</sub>MoO<sub>4</sub>. Zhur. neorg. khim. 9 no.11:2678-2679

N '64. (MIRA 18:1)

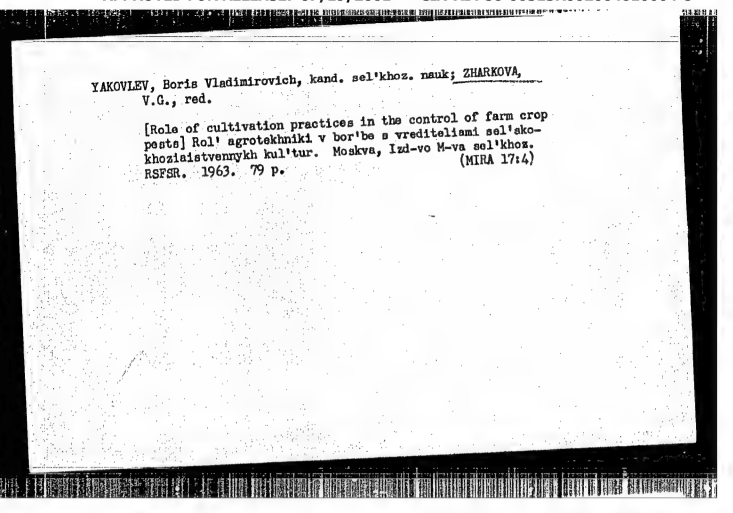
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M.V. Lomonosova.

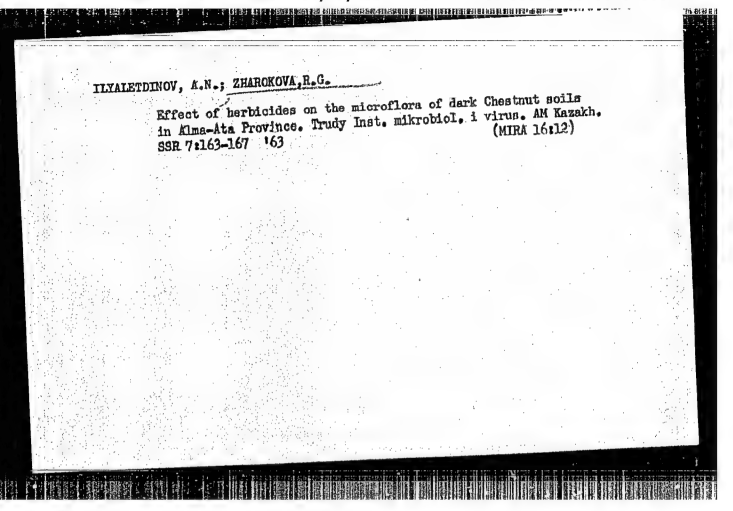
SKIOKIN, N.F.; ZHARKOVA, V.A.; MAKHON'KO, Yu.A.; SHAPOVALOV, Yu.S.

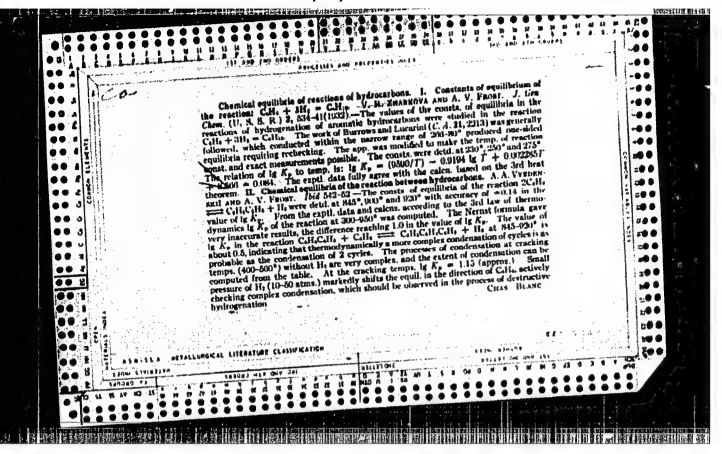
Prospective consolidated standards for the consumption of raw materials, fuel, electric power, and other materials in the production of ferrous metals in the U.S.S.R. Shor. trud. TSNIICHM no.45:172-181 '65.

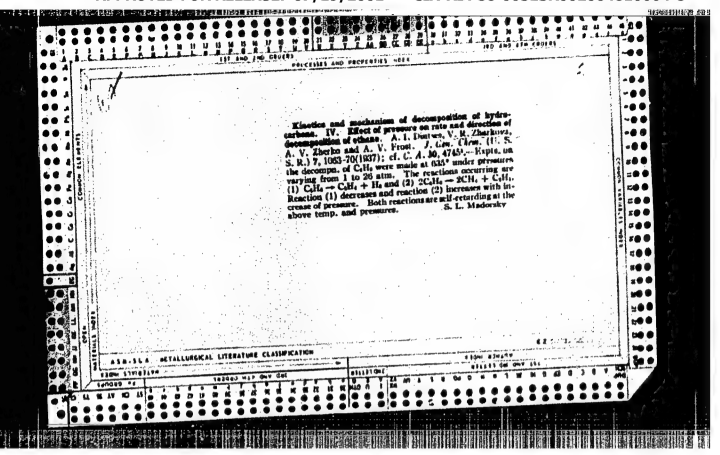
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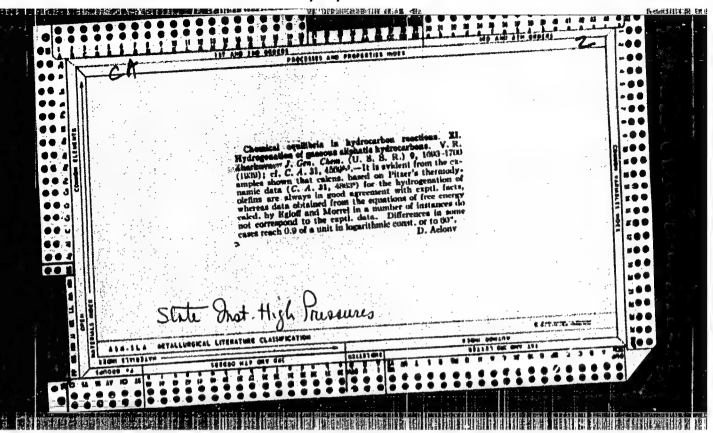
Methods of determining the volume and the composition of the U.S.S.R. metal stock by Union Republics and economic regions. Shor. trud. TSNIICHM no.45257-67 '65. (MIRA 18:9)

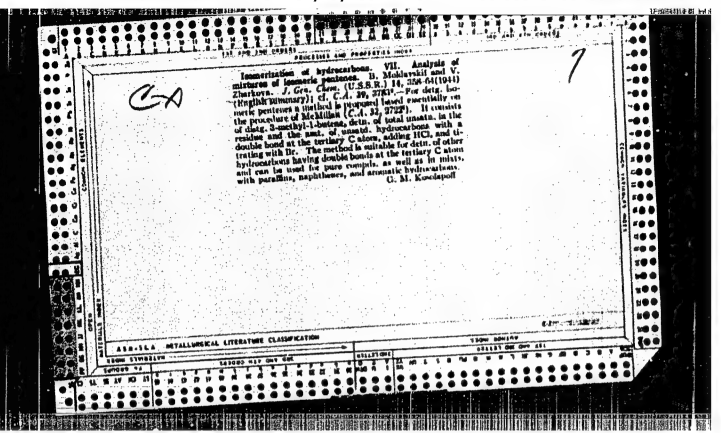


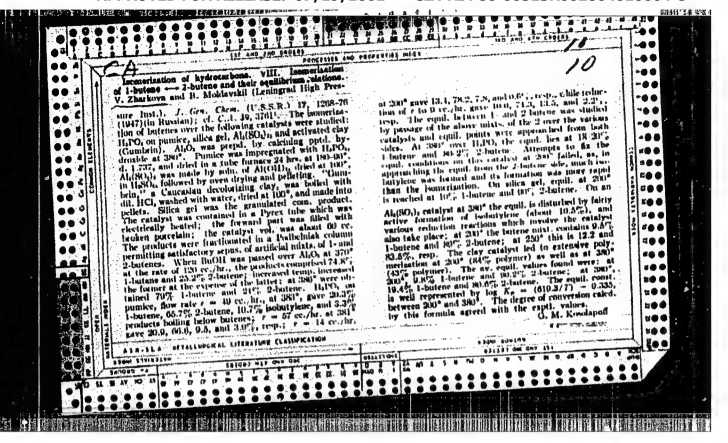












Methylation of hydrocarbons of the olefin series. B. L. Moldavskii, T. V. Nizovkina, and V. R. Zharkova (Leningrad High Pres. Inst.). J. Gen Chem (U.S.S.R.) 16, 127-34(1946)—In a study of El'tekov's methylation reaction (J. Russ. Phys. Chem. Soc. 10, 86(1878) of olefins by Mel in the presence of PbO, it was found that PbO causes side reactions of oxidative type with reduction of the oxide to Pb. It was also found that MgO may be substituted for PbO and MeCl for Mel. 2-Methyl-2-butene (60 g.) 51 g. MeCl. and 120g. PbO were heated in an autoclave 8 hrs. at 280-90°; this resulted in the formation of 55 g. Pb and 26g. PbCl2, while the org. products were composed of 65g. amylenes, 11.6g. hexeneheptenes,, and 16% diamylenes. Distn. through a Podbielniak column indicated the presence of isomeric amylenes in the 1st fraction, due to isomerization, while in thehigher fractions there were indications of the presence of 2,3-dimethyl-2butene, b. 730; 2,3,3-trimethyl-1-butene, b. 78°; and 2,3-dimethyl-1-butene, b. 55-8°. Repetition of the expt. at 265° for 10 hrs., using 71 g. 2-methyl-2butene, 252 g. MeCl, and 80 g. MgO, gave similarly 23% amylenes, 27.5% hexenes, and 31% heptylenes, with 14.3g. undistd. residue; fractionation of the products gave the same distribution as above. Use of mixed butenes instead of 2-methyl-2-butene (the mixt. used consisted of 1-butene 25%, 2-butene 6%, and isobutylene 6%), gave 54.5% C5-C7, olefins, which contained 3-methyl-1-butene and 2-methyl-2-butene, besides the products listed above; quant. G. M. Kosolanoff sepn. was not performed.

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PROST, Andrey Vladimirovich, prof. [deceased]. Prinimali uchastiye:

BUSHMAKIN, I.N.; VVEDENSKIY, A.A.; GRYAZNOV, V.M.; DEMENT'YEYA,

M.I.: DINTSES, A.I.; DOBRONRAVOV, R.K.; ZHARKQVA, V.R.; ZHERKO,

A.V.: IPAT'YEV, V.N.; KYYATKOYSKIY, D.A.; KOROBOV, V.V.; MOCR,

V.G.; NEMTSQV, M.S.; RAKOVSKIY, A.V.; REMIZ, Ye.K.; RUDKOVSKIY,

D.M.; RYSAKOV, M.V.; SERHERYAKOVA, Ye.K.; STEPUKHOVICH, A.D.;

STRIGALEVA, N.V.; TATEVSKIY, V.M.; TILICHEYEV, N.D.; TRIFEL',

A.G.: FROST, O.I.; SHILYAYEVA, L.V.; SHCHEKIN, V.V., DOLGOPOLOV,

N.M., SOSTAVITE! GERASIMOV, YA.I., etv.red.; SMIRNOVA, I.V.; red.;

TOPCHIYEVA, K.V.; YASTREBOV, V.V., red.; KONDRASHKOVA, S.F., red.

isd-va; LAZAREVA, L.V., tekhn.red.

[Selected scientific works] Isbrannye nauchnye trudy. Moskva, (MIRA 13:5)

1. Chlen-korrespondent AN SSSR (for Gerasimov). (Chemistry, Physical and theoretical)

Existing in fight of the control of

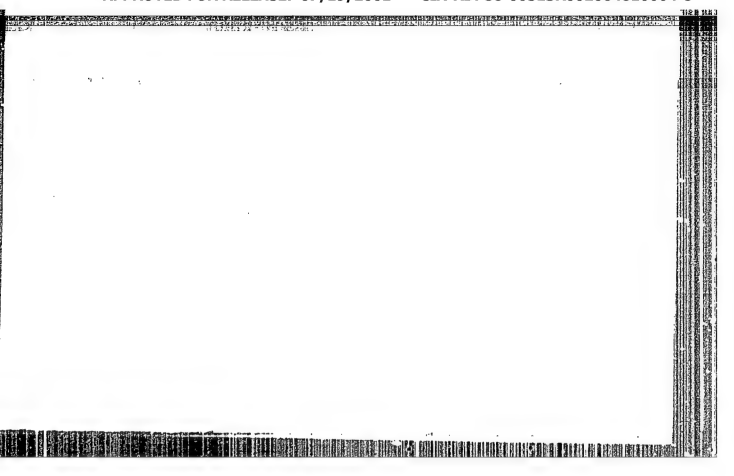
ZHARKOVA, Yu.V.; SADIKOV, B.A.; GORBATOV, Yu.B., retsenzent; SHCHERBAKOVA, Ye.A., retsenzent

[Problems in physics] Zadachnik po fizike. Moskva, Pt.2. 1963. 130 p. (MIRA 18:2)

1. Moscow. Energeticheskiy institut. 2. Kafedra fiziki Moskovskogo energeticheskogo instituta (for Gorbatov, Shcherbakova).

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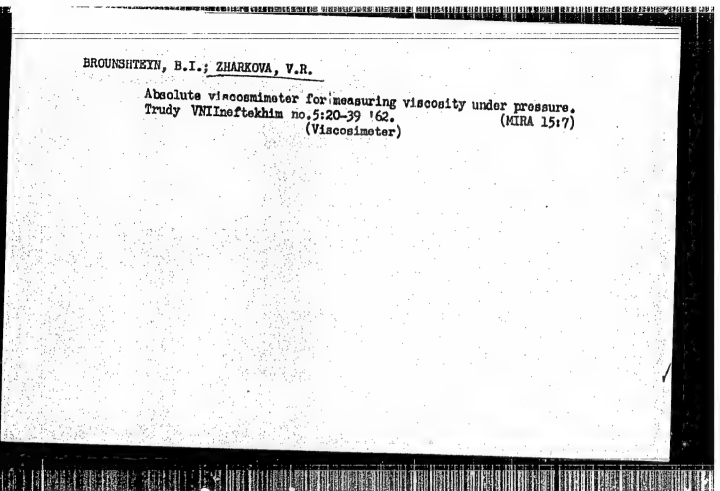


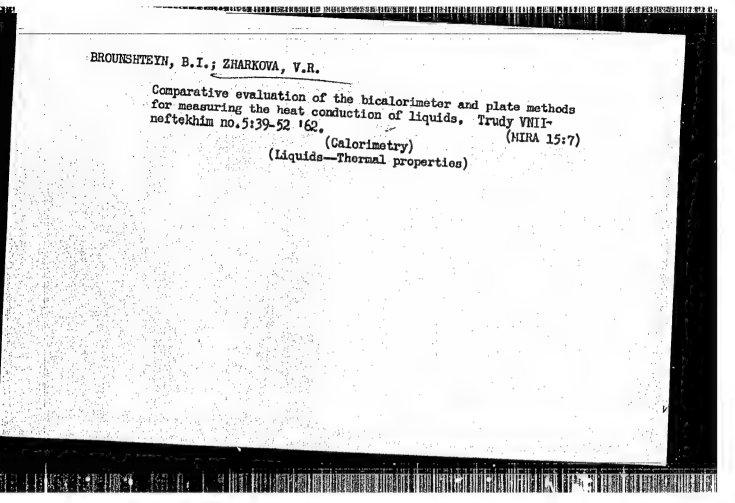


ORLOV, N.V.; ZHARKOVA, A.K.

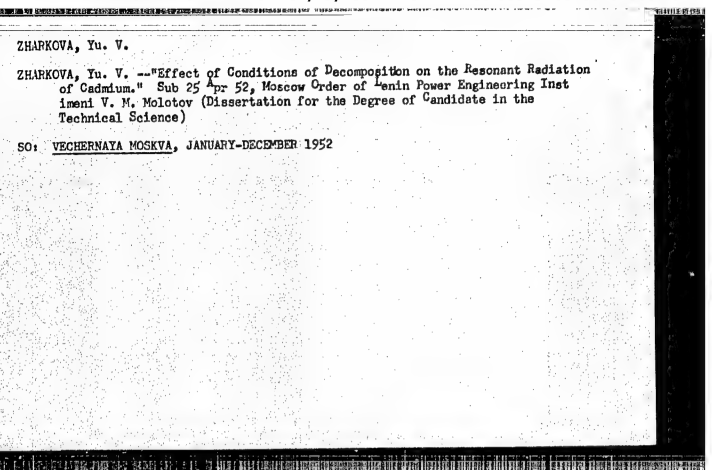
Fangotherapy in compound treatment of inflammatory diseases of the spinal cord. Vop. kur., fizioter. i lech. fiz. kul't. 29 no.1:3-6 '64. (MIRA 17:9)

1. Nervnoye otdeleniye (nauchnyy rukovoditel' - prof. D.G. Shefer) Sverdlovakogo instituta kurortologii i fizicterapii (dir.- kand. med. nauk N.V. Orlov).





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	ZIIARKOVA,	vv. V.	Alecharge to ance of Prof 51 by Acad G	TESSE / Physics - Reson (Co Resonance Radiation Papors, Yu. Y. Zhan Union Elec Eng Inst "Dok Ak Hank SSSR" Investigates the in of cadmium as a fun current strengths; pressure on electron and potential s for various current	
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L 3275L-66 EWP(1)/EWT(m)/T LJP(c) SOURCE CODE: UR/0190/66/008/004/0569 AP6012706 ACC' NRI AUTHOR: Zharikova, Z. F.; Reztsova, Ye. V.; Berestneva, Z. Ya.; Kargin, Physicochemical Institute im. L. Ya. Karpov (Fiziko-khimicheskiy institut) TITLE: The effect of supramolecular structure in rubbers on their mechanical properties SOURCE: Vysokomolekulyarnyye soyedineniya, v. 8, no. 4, 1966, 569-572 TOPIC TAGS: natural rubber, synthetic rubber, vulcanization, molecular structure ABSTRACT: The dependence of the mechanical properties of structures in thinram vulcanizates with natural rubber and synthetic polyisoprene, polybutadiene, and sodium butadiene rubbers, on its supramolecular structures was investigated. Vulcanized rubber with more ordered structure was found to possess superior mechanical properties. Change in mixing temperature (in the range of 25-70C) does not significantly affect the structure and properties of the rubber. Structure formation in thiuram polyisopreme vulcanized rubber subjected to stretching was investigated by electron microscopy. Ribbon-like structures were found to be perpendicular to the applied force during stretching of vulcanized rubbers. Orig. art. has: 4 fig-[BT] ures and 1 table. SUB CODE: 11/ SUBM DATE: 05Feb65/ ORIG REF: 007/ 678.03 第3+678.43 UDC:

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SOURCE CODE: UR/0053/66/088/003/0419/0438

AUTHOR: Zharkov, G. F.

ORG: Physics Institute im. P. N. Lebedev, AN SSSR (Fizicheskiy institut AN SSSR)

TITLE: The Josephson tunneling effect in superconductors

SOURCE: Uspekhi fizicheskikh nauk, v. 88, no. 3, 1966, 419-438

TOPIC TAGS: superconductivity, tunnel effect, wave function, tunnel current, nuclear binding energy

ABSTRACT: This is a review paper dealing the the tunneling of coupled electron pairs from the ground state (Fermi surface) of one superconductor to the Fermi surface of another through a thin insulating layer, first reported by B. Josephson (Phys. Letts. v. 1, 251, 1962). The treatment in the article is restricted to bulk superconductors. Subjects treated are the general nature of the Josephson effect, a phenomenological description of the effect and the role played by the wave function of the superconductor in this description, derivation of the equations for the relation between the Josephson current and the binding energy, the influence of the presence of magnetic and electric fields, the influence of the transverse dimensions of the barrier, the Josephson current in barriers connected

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in parallel and other interference effects, and the nonstationary Josephson effect. In addition to reviewing briefly the contents of many published and unpublished papers dealing with the foregoing topics, mentions is made of papers dealing with other aspects such as the interpretation of the Josephson effect from the point of view of the Ginzburg-Landau theory, of the quasi-spin approach or of paramagnetic impurities in the superconductor, and some possible practical applications of the effect. Orig. art. has: 14 figures and 39 formulas.

SUB CODE: 20/ SUEM DATE: 00/ ORIG REF: 017/ OTH REF: 046

Card 2/2 CC

Translation from: Referativnyy zhurnal, Metallurgiya, 1957, Nr 6, p 257 (USSR)

AUTHORS: Zharkova, Z.P., Zhacheva, Ye.I.

TITLE: A Method for the

A Method for the Analytical Precipitation of Gold With Hydrazine Hydrochloride in Gold Electrolytes (Analiticheskiy metod osazhdeniya zolota solyanokislym gidrazinom v zolotykh elektrolitakh)

PERIODICAL: Nauch.-issled. tr. Tsentr. n.-i. in-t vspomogat. izdeliy i zapas. detaley k tekstil'n. oborud., 1956, Nr 4, pp 44-47

ABSTRACT: A 10-cc test sample is taken for the determination of Au in cyanide electrolytes that contain 3-4 g/l of the metal. For concentrated to 500 cc with water and a 20-50 cc aliquot is used for the analysis. The test sample is neutralized with HCl to phenolphthalein and heated boiling. 50 cc of 10% hydrazine hydrochloride are added to the boiling solution and the boiling is continued to the complete reduction Au according to the following reaction:  $4KAu(CN)_2 + N_2H_4 \cdot 2HCl = 4Au + N_2 + 2KCl + 6HCN + 2KCN$ . The brown Au precipitate is filtered off and calcined in a porcelain crucible at 800°C. The Au in the

A Method for the Analytical Precipitation of Gold (cont.)

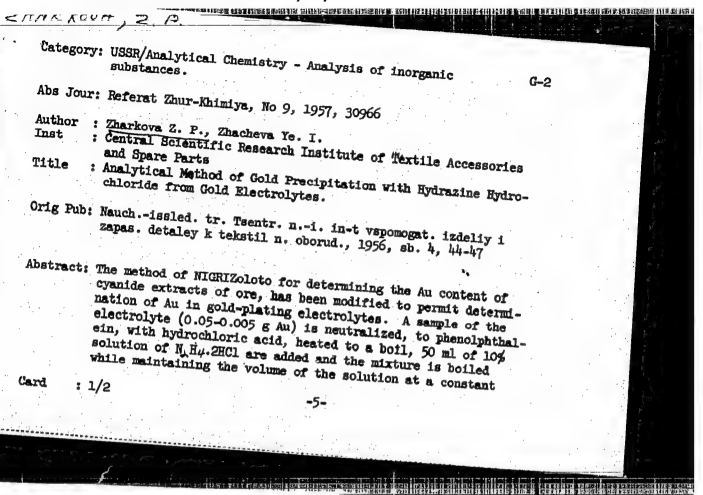
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crucible is dissolved in 5 cc aqua regia and evaporated three times on a water bath with 1:3 HCl. Then 3 cc of 1:50 HCl and 2-3 drops of Br water are added to oxidize the monovalent Au. The contents of the crucible are evaporated to 1.5 total volume (sic!). The remaining solution is filtered, the paper is washed until the added. 2-3 drops of orthodianizidine solution are then added to the filtrate which the appearance of a light blue color of dispersed Au. The Au content of the solution is calculated according to the following reaction:

2AuCl<sub>3</sub>+3C<sub>6</sub>H<sub>6</sub>O<sub>2</sub>=2Au+3C<sub>6</sub>H<sub>4</sub>O<sub>2</sub>+6HCl.

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Category: USSR/Analytical Chemistry - Analysis of inorganic

substances.

Abs Jour: Referat Zhur-Khimiya, No 9, 1957, 30966

level. The Au precipitate is filtered off, washed with hot water, diried and calcined at 800°. To the filtrate are added 25 ml of 10% solution of N.H., 2HCl and the mixture is boiled to check completeness of precipitation. The calcined precipitate is dissolved in 5 ml of aqua regia, evaporated, and the residue is evaporated to dryness, 3 times, with 1-2 ml dilute HCl (1:3). 3 ml of dilute HCl (1:50) are added, followed by 2-3 drops of a hydrochloric acid solution of Br2, and the mixture is evaporated to dryness. The residue is diluted to 30-40 ml, combined with 0.1 g K-bifluoride, 2-3 drops of o-dianisidine solution (10 mg in 10 ml HCl, 1:300) and titrated, after 5 minutes, with a solution of hydroquinone (0.4180 g dissolved in 500 ml water, acidified with 10 ml concentrated HCl, and diluted with dilute hydrochloric acid, 1:300, at a ratio of 1:100) until a colorless or faintly bluish solution is obtained (2AuCi<sub>3</sub> + 3C<sub>6</sub>H<sub>6</sub>O<sub>2</sub>  $\longrightarrow$  3C<sub>6</sub>H<sub>6</sub>O<sub>2</sub>+ 2Au + 6HCl).

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